Structural Variants in the Soybean Genome Localize to Clusters of Biotic Stress-Response Genes


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Genome-wide structural and gene content variations are hypothesized to drive important phenotypic variation within a species. Structural and gene content variations were assessed among four soybean (Glycine max) genotypes using array hybridization and targeted resequencing. Many chromosomes exhibited relatively low rates of structural variation (SV) among genotypes. However, several regions exhibited both copy number and presence-absence variation, the most prominent found on chromosomes 3, 6, 7, 16, and 18. Interestingly, the regions most enriched for SV were specifically localized to gene-rich regions that harbor clustered multigene families. The most abundant classes of gene families associated with these regions were the nucleotide-binding and receptor-like protein classes, both of which are important for plant biotic defense. The colocalization of SV with plant defense response signal transduction pathways provides insight into the mechanisms of soybean resistance gene evolution and may inform the development of new approaches to resistance gene cloning.

Genetic variation within and between species is most commonly quantified by single nucleotide polymorphisms (SNPs). There has been increased interest in recent years to also resolve genetic differences in terms of structural variation (SV), which includes copy number variation (CNV) caused by large insertions and deletions, and other types of rearrangements such as inversions and translocations. The copy number of a specific gene or gene family has been associated with variation for specific traits, such as the digestion of starchy foods in humans (Perry et al., 2007), boron toxicity tolerance and winter hardiness in barley (Hordeum vulgare; Sutton et al., 2007; Knox et al., 2010), dwarfism and flowering time in wheat (Triticum aestivum; Pearce et al., 2011; Díaz et al., 2012), and insecticide and virus resistance in Drosophila melanogaster (Schmidt et al., 2010; Magwire et al., 2011). Genomic SV is thought to be an important factor in determining phenotypic variation for a wide range of traits (for review, see Stankiewicz and Lupski, 2010).

SV studies have been published in various invertebrate (Dopman and Hartl, 2007; Emerson et al., 2008; Maydan et al., 2010) and mammalian (Graubert et al., 2007; Guryev et al., 2008; Lee et al., 2008; Perry et al., 2008; Gazave et al., 2011; Golzio et al., 2012) systems; however, a high proportion of such studies have been conducted in humans, where there is interest in identifying associations between SV, complex diseases, and neurological disorders (Conrad et al., 2010; Craddock et al., 2010; Sudmant et al., 2010; Girirajan et al., 2011). Domesticated animal species, including dog, cow, and silkworm, have also been the focus of recent investigations of SV (Chen et al., 2009; Nicholas et al., 2009, 2011; Liu et al., 2010; Sakudoh et al., 2011; Clop et al., 2012). CNV have been identified within genes and gene families associated with specific biological functions, such as immunity. Some evidence from these studies suggests that phenotypic variation caused by CNV can rapidly emerge and be driven to fixation by breeders.

Recent studies in maize (Zea mays) have explored the exceptionally high rates of SV between inbred accessions (Springer et al., 2009; Beló et al., 2010; Swanson-Wagner et al., 2010). The maize profile indicates that there are continuously high levels of SV between
acrossions throughout all 10 chromosomes, interspersed by relatively small regions of conservation thought to be regions of identity by descent. Other than this work in maize and work in Arabidopsis (Arabidopsis thaliana; Santuari et al., 2010; Cao et al., 2011; Gan et al., 2011; Lu et al., 2012), rice (Oryza sativa; Yu et al., 2011; Xu et al., 2012), and sorghum (Sorghum bicolor; Zheng et al., 2011), relatively little is known about the intraspecific structural genomic variation within plant species. Maize is a primarily outcrossing species known to have a remarkably diverse germplasm (Wright et al., 2005; Gore et al., 2009); most other domesticated plant and crop species would be expected to have lower rates of SV as a consequence of their narrower genetic base. Soybean (Glycine max) is an interesting system for comparison. Soybean is a self-pollinating species with a comparatively narrow genetic base that has experienced severe genetic bottlenecks during domestication (Hyten et al., 2006). Soybean SNP rates among accessions are relatively low, typically on the order of one SNP per kb (Hyten et al., 2006; Lam et al., 2010). Given the low polymorphism rate and narrow genetic base, one might surmise that soybean accessions are similarly devoid of SV. However, a recent study using microarray-based comparative genomic hybridization (CGH) analysis found surprisingly high rates of SV between two cultivars (‘Kingwa’ and ‘Williams’; Haun et al., 2011). Additionally, regions of SV were identified within sublineages of the reference cultivar (‘Williams 82’), including several genetic loci that exhibited presence-absence variation (PAV) among the different Williams 82 individuals (Haun et al., 2011). PAV is a subclass of CNV in which a specific gene or other sequence is present in some accessions and entirely absent in others (Springer et al., 2009).

In this study, we sought to define the range of SV between four genetically diverse soybean accessions: ‘Archer’, ‘Minsoy’, ‘Noir 1’, and Williams 82. Minsoy and Noir 1 are plant introductions, while Archer is a North American cultivar. These three genotypes were of particular interest because they are the parental lines for the recombinant inbred line populations utilized in the first agronomic quantitative trait loci (QTL) analyses of soybean (Lark et al., 1995; Orf et al., 1999) and represent a wide range of soybean sequence diversity (Zhu et al., 2003). Zhu et al. (2003) reported that approximately two-thirds of the common soybean SNPs found in a set of 25 diverse lines are polymorphic among these three genotypes. Williams 82 is a modern soybean cultivar that provided the reference genome sequence (Schmutz et al., 2010). Coarse structural differences between the four genotypes (i.e. CNV) were resolved using CGH technology, and specific gene content variants (i.e. gene PAV) were identified using exome-resequencing analyses. These data were used to catalog the set of genes located within regions enriched for SV, giving new insight into the mechanisms and forces that may be driving SV in soybean.

RESULTS

SNP Variation among Four Soybean Genotypes

Little is known about the genomic SV among soybean accessions. Furthermore, the relationship between genomic SV and haplotype variation is essentially unstudied. To investigate this relationship, we generated whole-genome CGH data and comprehensive SNP genotype data on the four soybean genotypes Archer, Minsoy, Noir 1, and Williams 82. There is known to be genetic heterogeneity within some soybean cultivars (Fasoula and Boerma, 2007; Haun et al., 2011; Varala et al., 2011; Yates et al., 2012), meaning that there can be differences between individual plants or sublines within an accession. Therefore, the four genotypes in this study were each represented by a single plant. Herein, the four individuals are referred to by their abbreviated or cultivar name (Archer, Minsoy, Noir 1, and Wm82) for brevity and simplicity, but their full subline names are given in the Materials and Methods.

Comprehensive SNP genotyping data were generated using the Illumina Infinium platform for soybean, which consists of approximately 44,000 SNPs spaced at regular intervals across the soybean genome. The SNP profiles of the Archer, Minsoy, and Noir 1 individuals were compared with the Wm82 individual (Fig. 1). The three genotypes all displayed discontinuous patterns of polymorphism along the 20 chromosomes. Archer showed the lowest level of polymorphism relative to Wm82, including several stretches that appeared to be shared haplotypes across long chromosomal regions (e.g. chromosomes 3, 19, and 20 in Fig. 1). These shared haplotypes may be regions of identity by descent, as Williams 82 was the Phytophthora root rot resistance donor (Rps1k) in the Archer pedigree (Cianzio et al., 1991). Minsoy and Noir 1 also appeared to have some haplotype regions shared with Wm82, although to a lesser degree than Archer (Fig. 1).

Genomic SV among Four Soybean Genotypes

To gauge SV between genotypes, Archer, Minsoy, and Noir 1 were each hybridized with Wm82 as the reference in CGH experiments (Supplemental Fig. S1). Among the three comparisons, the number of genomic segments exhibiting significant CNV ranged from 188 to 267 segments per genotype comparison (Table I). The CNV false discovery rate based on technical variables is likely low, as a control Wm82-Wm82 self-hybridization identified only 13 significant CNV (Table I).

The median size of a CNV segment was approximately 18 to 23 kb for all three comparisons (Table I). The distribution of significant CNV was discontinuous throughout each comparison (Supplemental Fig. S1). The chromosomes exhibited differing levels of SV, including whole chromosomes with little to no evidence of SV (e.g. chromosomes 5 and 11) and chromosomes...
with extended regions of SV (e.g. chromosomes 3 and 18; Fig. 2).

Hundreds of genes colocalized to regions of SV. The gene models overlapping with significant CNV regions were identified for each genotype comparison (Supplemental Table S1). In total, 672 different gene models overlapped with CNV regions in at least one of the three comparisons (Supplemental Table S1). Over one-third (37%) of the 672 gene models were significant in more than one comparison, and 120 were significant in all three comparisons (Supplemental Fig. S2). The false discovery rate is likely to be low, as the control Wm82-Wm82 self-hybridization identified only one gene model within a significant CNV. Furthermore, CGH technical replications of Minsoy-Wm82 and Noir 1-Wm82 were performed to estimate the reproducibility of these discoveries. While the technical replicate hybridizations had more noise than the original hybridizations, the CNV patterns were essentially the same (Supplemental Fig. S3). Importantly, the vast majority of gene models identified within the CNV regions in the technical replicates were also significant in the original hybridizations (77% for Minsoy-Wm82 and 77% for Noir 1-Wm82; Supplemental Table S1).

Most of the polymorphic loci consist of DownCNV (Table I), which can be interpreted as regions in which the tested genotype has fewer DNA copies than the reference genotype Wm82 or that hybridize less efficiently than Wm82 (presumably due to nucleotide sequence polymorphisms). The DownCNV can be visualized as downward peaks in Figure 2 and Supplemental Figure S1. The high frequency of DownCNV relative to UpCNV is expected, as the microarray was developed based on the Williams 82 reference genome sequence (Schmutz et al., 2010). Significant UpCNV were observed in some instances (approximately 12% of all CNV; Table I), most prominently along chromosomes 3 and 7. These UpCNV occur within regions of known intracultivar heterogeneity, in which the Williams 82 reference genome sequence does not perfectly match the Wm82-ISU-01 haplotype (Haun et al., 2011). The UpCNV likely represent genomic regions that are absent from Wm82-ISU-01, present in the Williams 82 reference sequence, and also present in the respective genotypes (Archer, Minsoy, and/or Noir 1) in which the UpCNV is observed. There are also some examples of UpCNV that do not occur within regions of known heterogeneity (e.g. the UpCNV on chromosome 12) and that possibly represent copy number gains in the respective genotypes relative to the Williams 82 genome.

The relationship between the genomic SV profiles and the Infinium SNP variation profiles is shown in Supplemental Figure S4. The interpretation of this comparison is complicated because the Infinium SNP assays were selected by virtue of being highly polymorphic, while the CGH probes were selected without regard to their potential to underlie CNV. This ascertainment bias leads to a distortion in the fraction of data that the two technologies detect as polymorphic. Despite this complication, some general trends were observed. SNP and SV rates were generally coincident along each chromosome, such that regions of high SV were localized to regions with high rates of nucleotide polymorphism. The colocalization of SNP and CNV suggests that areas rich for both SNP and SV represent divergent haplotypes for a given comparison. However, exceptional instances were observed in which regions with profound SNP polymorphism rates exhibited little SV (e.g. Archer-Wm82 chromosome 1) or regions with abundant CNV exhibited little to no SNP variation (e.g. Archer-Wm82 chromosome 3; Supplemental Fig. S4).

Exome capture and resequencing data were generated for the four soybean genotypes to validate the array CGH data and further evaluate gene content variation between the lines using a fixed exon content

Figure 1. SNP genotyping reveals regions of conservation and divergence between Wm82 and Archer (A), Wm82 and Minsoy (B), and Wm82 and Noir 1 (C). Gray spots indicate matching SNPs, and black spots indicate polymorphic SNPs between genotypes. The spots are jittered along the x axis to enhance the resolution of the data points.
measure. A stringent analysis pipeline was developed to identify gene content variation among the four genotypes based on the number of normalized exome read counts mapping to each gene model (for a description of the analysis details, see “Materials and Methods”). This allowed for the identification of a subset of genes that exhibited high read counts (more than 30) in at least one genotype and zero read counts in at least one other genotype. These 133 genes make up the high-confidence list of PAV. The locations of the PAV along each chromosome are shown as red spots in Figure 2 and Supplemental Figure S1. The gene models and presence-absence profiles of these 133 genes are shown in Supplemental Table S2, and the distribution of “absent” genes among the four genotypes is shown in Supplemental Figure S5.

The CGH and exome-resequencing data were compared to cross-validate the PAV calls on the 133 genes. The log2 ratios of the exome-resequencing counts and CGH data for each gene × genotype comparison are shown in Figure 3. There was a strong significant correlation (P < 0.0001) between the CGH and exome-resequencing platforms for all of the genotype comparisons. Furthermore, the majority (58%) of the PAV genes called absent in Archer, Minsoy, or Noir 1 relative to Wm82 were located within significant CNV segments.

The 133 PAV genes identified represent a high-confidence list but almost certainly underestimate the number of genes that have full or partial gene content variation among the tested genotypes (for further explanation, see Supplemental Materials and Methods S1). Additionally, any novel gains in gene content exhibited by the lines would be missing from this analysis because the exon capture can only deliver sequences homologous to the probes designed from the Williams 82 reference sequence.

PCR assays were conducted to estimate the presence-absence distribution of high-confidence PAV genes in a sample of 31 germplasm accessions, including 10 modern cultivars, 17 North American ancestors, and four landraces. PCR primers were designed within the predicted coding regions of 13 putative PAV genes (Supplemental Table S3); these 13 genes also showed evidence of CNV in the CGH experiment. The presence-absence state of these 13 genes was tested by PCR on the full set of 31 accessions (Supplemental Table S4; Supplemental Fig. S6). A wide range of gene content variation among the accessions was observed for the two genes. Two genes, Glyma05g24800 and Glyma17g18890, showed present amplicons in all of the accessions except for the genotype in which absence was initially observed (Archer and Noir 1, respectively). Thus, these two genes are likely to be rare variants in which the absent state is found in few genotypes. The remaining 11 genes exhibited absent rates ranging from 13% to 65% of the accessions (Supplemental Table S5).

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Segments</th>
<th>UpCNV</th>
<th>DownCNV</th>
<th>Mean Size (bp)</th>
<th>Median Size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Archer versus Wm82</td>
<td>188</td>
<td>34</td>
<td>154</td>
<td>62,287</td>
<td>18,433</td>
</tr>
<tr>
<td>Minsoy versus Wm82</td>
<td>232</td>
<td>12</td>
<td>220</td>
<td>66,298</td>
<td>22,776</td>
</tr>
<tr>
<td>Noir 1 versus Wm82</td>
<td>267</td>
<td>29</td>
<td>238</td>
<td>54,499</td>
<td>18,733</td>
</tr>
<tr>
<td>Wm82 versus Wm82</td>
<td>13</td>
<td>11</td>
<td>2</td>
<td>11,017</td>
<td>7,679</td>
</tr>
</tbody>
</table>

Table 1. Summary statistics of soybean CNV number and size

Figure 2. CNV among soybean genotypes on chromosomes 3 (A) and 18 (B). Log2 ratios between each genotype relative to the Wm82 reference are shown. Blue spots indicate probes within significant CNV segments with values beyond threshold. Red spots indicate probes within PAV genes as determined by exome-resequencing analysis. The plots at the bottom of both A and B indicate the average variance (along a sliding window of 250 probes) between the log2 ratios of the Archer, Minsoy, and Noir 1 hybridizations.
High Levels of SV Associated with Disease Resistance Genes

The full set of genes associated with CNV or PAV regions, along with their Gene Ontology (GO) annotations and protein family prediction, is shown in Supplemental Table S5. GO analyses revealed 24 categories that were significantly enriched and nine categories that were significantly depleted for genes within CNV regions relative to all genes not found within significant CNV regions (Fisher’s exact test; multiple testing adjusted $P < 0.05$; Supplemental Table S6). Although the level of significance varied for each genotypic comparison, the direction (enrichment or depletion) was consistent among the Archer, Minsoy, and Noir 1 comparisons with Wm82 (Supplemental Table S6).

Genes within regions of structural and gene content variation frequently had potential functions in disease resistance and response to biotic stress. GO categories with the greatest enrichment of genes in CNV regions were related to plant-pathogen interactions and included “defense response,” “plant-type hypersensitive response,” “programmed cell death,” and “apoptosis.” A specific enrichment was observed for genes encoding nucleotide-binding (NB) proteins and receptor-like proteins (RLPs; Fisher’s exact test; $P_1 = 1.87 \times 10^{-58}$ and $P_2 = 4.32 \times 10^{-122}$, respectively; Table II), which often function in disease resistance (Krujit et al., 2005; DeYoung and Innes, 2006). Enrichment of NB- and RLP-encoding genes was also observed in PAV (Fisher’s exact test; $P < 0.01$; Supplemental Table S6). The full list and genomic positions of soybean gene models identified as NB (392 gene models) and RLP (220 gene models) are shown in Supplemental Tables S7 and S8, respectively. Figure 4 shows the colocalization of these gene classes with CNV spikes. The hybridization variances among the Archer-Wm82, Minsoy-Wm82, and Noir 1-Wm82 comparisons are shown, revealing the approximate locations of structurally varying genes.

<table>
<thead>
<tr>
<th>Genes</th>
<th>All Eligible Genes</th>
<th>NB Encoding</th>
<th>RLP Encoding</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>CNV</td>
<td>%CNV</td>
</tr>
<tr>
<td>Unique genes</td>
<td>10,096</td>
<td>188**</td>
<td>1.9</td>
</tr>
<tr>
<td>Small multigene family (2–10)</td>
<td>28,299</td>
<td>247**</td>
<td>0.9</td>
</tr>
<tr>
<td>Large multigene family (more than 10)</td>
<td>7,858</td>
<td>237**</td>
<td>3.0</td>
</tr>
<tr>
<td>Isolated multigene family members</td>
<td>30,928</td>
<td>196**</td>
<td>0.6</td>
</tr>
<tr>
<td>Clustered multigene family members</td>
<td>5,229</td>
<td>288**</td>
<td>5.5</td>
</tr>
<tr>
<td>Genes containing tandem repeats</td>
<td>13,684</td>
<td>217</td>
<td>1.6</td>
</tr>
<tr>
<td>Genes with nearby TEs</td>
<td>11,347</td>
<td>250**</td>
<td>2.2</td>
</tr>
<tr>
<td>All CNV-eligible genes</td>
<td>46,253</td>
<td>672</td>
<td>1.5</td>
</tr>
</tbody>
</table>

*%CNV is calculated as the number of CNV in a particular category divided by the total number of genes in that category. **Fold change is relative to the genome-wide %CNV: the %CNV for a particular category divided by genome-wide %CNV (1.5%).

Figure 3. Cross-validation of CGH and exome-resequencing data for the 133 genes identified as PAV in the exome-resequencing data set. Archer/Wm82 (A), Minsoy/Wm82 (B), and Noir 1/Wm82 (C) log2 ratios based on CGH (x axis) and exome-resequencing counts (y axis) each exhibited significant correlations. Each spot represents one of the 133 genes. Spot coloration is based on the gene presence/absence call in the exome-resequencing data. Red spots indicate genes called absent in Archer (A), Minsoy (B), or Noir 1 (C). Blue spots indicate genes called present in the respective genotype and absent in Wm82. Gray spots indicate genes called present in both the respective genotype and Wm82. Black spots indicate genes called absent in both the respective genotype and Wm82.
conserved regions (shown as relatively flat intervals) as well as structurally diverged regions (shown as peaks or peak clusters) among the Archer, Minsoy, and Noir 1 genomes. The locations of all annotated NB (red) and RLP (blue) genes in the genome.

The locations of structurally diverged regions were compared with the results of previous soybean genetic mapping studies, which are publicly available at http://www.soybase.org. The regions with the greatest amplitude of CNV variance account for 94 centimorgans of the soybean composite genetic map, which is less than 4% of the total map. Previous genetic mapping experiments in soybean indicate that multiple QTL and/or genes for disease resistance map to nearly all of the regions with highest CNV variance (genetic mapping data obtained from http://www.soybase.org). Fourteen percent (43 of 311) of QTL for disease resistance map to the regions with highest CNV variance, while only 7% (85 of 1,221) of non-disease-related QTL map to these regions. This suggests that a large portion of the qualitative and quantitative variation in disease resistance may be derived from gene content variation in NB- or RLP-encoding gene clusters.

The physical arrangement, copy number, and repetitive nature of genes residing within regions of CNV were assessed to determine whether the enrichment of the NB- and RLP-encoding genes may be influenced by factors other than their functions in disease resistance and the selection pressures potentially associated with those functions (Bishop et al., 2000; Mondragón-Palomino et al., 2002; Chen et al., 2010). Gene family size appeared to be one such factor; CNV regions were associated with large gene families more often than predicted by random expectations (Table II; Supplemental Table S9). Seventeen percent of predicted genes in the soybean genome are members of large multigene families (families with more than 10 members). In comparison, 35% of genes within CNV regions in our study were members of large multigene families (Fig. 5A).
The relationship between large gene families and CNV regions appears to be primarily due to the NB- and RLP-encoding gene families (Fig. 5A). When these genes, representing 4% of the large multigene families, 0.5% of the small multigene families, and 1% of the unique genes, are removed from the analysis, the frequency of large gene family members within CNV is near the frequency that would be expected at random. These correlations indicate that CNV occur more frequently within the NB- and RLP-encoding families than other large multigene families. It is evident that not all large gene families are associated with CNV regions and that there are requirements for CNV beyond gene family size.

The physical distribution of the members of multigene families appears to be a key component of the SV-enriched regions. Only members of multigene families that are located within clusters tend to be associated with CNV regions (Fig. 5B); isolated family members that are not clustered do not often associate with CNV regions. This trend is observed across all classes of genes (Fig. 5B); the tightly clustered NB- and/or RLP-encoding families on chromosomes 3, 6, 7, 16, and 18 are notable examples (Fig. 4).

Genes that contain tandem repeats within CNV regions were identified at a slightly higher frequency than expected, with 30% of all genes containing tandem repeats and 32% of genes with CNV containing tandem repeats (Fig. 5C; Supplemental Table S10). In comparison with all genes, there is a slightly higher percentage of tandem repeats in the NB and RLP classes (32% and 39%, respectively). In addition, there is a higher percentage of tandem repeats in both NB- and RLP-encoding genes within CNV regions (37% and 55%, respectively; Fig. 5C). This association of tandem repeats to genes within CNV is specific to the NB and RLP classes and is not observed in the remainder of the genic sequences (Fig. 5C), indicating that the presence of tandem repeats without other factors attributed to NB- and RLP-encoding genes does not influence CNV.

Transposable elements (TEs) can result in duplication, deletion, or transposition of nearby non-TE genes through a variety of mechanisms (Bennetzen, 2000). Possible evidence of TE-mediated transposition and deletion of NB-leucine-rich repeat (LRR)-encoding genes in soybean has been reported previously (Innes et al., 2008; Wawrzynski et al., 2008). We observed an enrichment of nearby TEs (Du et al., 2010) in genes located within CNV regions in comparison with genes not located within CNV regions (Fisher’s exact test; \( P < 0.05 \); Fig. 5D). However, NB- and RLP-encoding genes within CNV regions have no significant enrichment for nearby TEs (Table II; Fig. 5D). These observations indicate that while TEs may play an important role in generating CNV, the CNV regions surrounding NB- and RLP-encoding genes are either primarily generated by other mechanisms or nearby TEs were sufficiently fragmented to not be detectable by methods implemented to identify partial TEs for inclusion in SoyTEdb (Du et al., 2010).
Confirmation of CGH Trends on Additional Soybean Accessions

We performed additional CGH experiments with additional soybean accessions to test whether the trends observed in the CGH experiments involving Archer, Minsoy, Noir 1, and Wm82 would also be observed in other genotype comparisons. The first two hybridizations utilized Wm82 as the reference dye and genotypes 'Essex' and 'Richland' as the experimental dye. Essex is a cultivar release from 1973 (Smith and Camper, 1973), and Richland is a North American ancestor accession. A third hybridization directly compared the cultivars Archer and ‘M92-220’, which is a recently developed cultivar derived from the 2006 Crop Improvement Association seed stock of cv ‘MN1302’ (Orf and Denny, 2004). The last hybridization used for comparison involved accessions Kingwa and Williams. This hybridization was described previously (Haun et al., 2011) but was never analyzed for the patterns of CNV regional abundance.

The CNV frequency and size for each of the hybridizations are shown in Supplemental Table S11. The gene models associated with significant CNV are shown in Supplemental Table S12, and the enriched GO and Pfam categories are shown in Supplemental Table S13. The patterns and trends observed in these additional hybridizations closely paralleled the results of the Archer, Minsoy, Noir 1, and Wm82 experiments. For each of the four additional hybridizations, defense-related GO terms and Pfam domains associated with plant resistance genes had significant overrepresentation within genes in CNV regions (Supplemental Table S13). Combining the Essex-Wm82 and Richland-Wm82 data with the previous experiments on Archer, Noir 1, and Minsoy, 782 different gene models overlapped with CNV regions in at least one of the five comparisons (Tables S1 and S12). Nearly half (46%) of these gene models were significant in more than one comparison (Supplemental Fig. S7).

DISCUSSION

Distribution and Rates of SV in the Soybean Genome

The CNV profiles in this study indicate that soybean has relatively long chromosomal regions (and nearly entire chromosomes) that exhibit virtually no SV among genotypes, interspersed with pockets of high SV ranging from several kb to greater than 10 Mb in length. DownCNV (segment loss relative to Wm82) were much more abundant than UpCNV (segment gain relative to Wm82). This is expected, considering that the reference dye in all of the hybridizations was Wm82, which serves as the reference sequence used to design the microarray platform. The relative abundance of DownCNV is consistent with previous CGH studies of similar design, including those of maize (Springer et al., 2009; Swanson-Wagner et al., 2010) and rice (Yu et al., 2011). The patterning of the statistically significant DownCNV resembles that observed in soybean fast-neutron deletion lines (Bolon et al., 2011), indicating that the most prominent peaks likely represent missing genomic regions within the Archer, Minsoy, and/or Noir 1 genotypes relative to Wm82. The rare UpCNV likely represent heterogeneous regions within the Williams 82 cultivar (Haun et al., 2011) and may represent sequences and gene content variants absent in the particular Wm82 individual used in these hybridization experiments. Collectively, these findings indicate that many of the SV detected in this study are likely caused by DNA segments that are present in some lines and absent in others.

Conversely, nucleotide polymorphism among lines may have also contributed to the microarray hybridization differences. This is particularly likely within large segments, in which true SV may be interspersed with regions of high sequence polymorphism. For example, the largest CNV identified in this study was a nearly 2-Mb DownCNV located in a gene-poor pericentromeric region of chromosome 4. This region, which is approximately three times larger than the next largest CNV identified in this study, was barely beyond the significance threshold in the Archer-Wm82 and Minsoy-Wm82 comparisons. Clearly, this region is not a true 2-Mb deletion in Archer and Minsoy but instead may indicate a combination of SV and SNP polymorphisms throughout the region. In this sense, the CGH analysis represents a scan of genome-wide polymorphisms that is particularly sensitive to identifying strong SV (but the “CNV” terminology is not necessarily an accurate description for all of the polymorphic segments identified in the analyses). However, the low SNP rates in soybean (Hyten et al., 2006) and the application of stringent significance thresholds provide confidence that a sizeable fraction of polymorphic segments identified in this study consists primarily of true SV. A PCR survey of 31 accessions supported this conclusion, indicating that the subset of CNV and PAV identified in this study likely represented a range of both rare and common structural variants throughout the soybean germplasm.

The most extensive studies of crop plant SV to date have been performed in maize (Springer et al., 2009; Swanson-Wagner et al., 2010). In terms of CNV distribution, the soybean comparisons are virtually the opposite of what has been observed in maize. In maize, accessions tend to exhibit high rates of SV throughout their chromosomes, with infrequent regions of structural conservation interspersed. In soybean, chromosomes tend to exhibit long stretches of conservation interspersed with regions enriched for CNV. The relative rates of SV between the two species are consistent with published rates of nucleotide variation, in which domesticated maize lines exhibit much higher SNP rates than domesticated soybean lines (Wright et al., 2005; Hyten et al., 2006; Gore et al., 2009; Lam et al., 2010). Not surprisingly, the rates of SV between soybean genotypes are also an order of magnitude lower that that of domesticated maize.
magnitude less than the differences observed between the genome sequences of soybean and its nearest wild relative *Glycine soja*, in which more than 1,000 genes are estimated to have large structural differences caused by deletions, insertions, inversions, transpositions, or translocations (Kim et al., 2010).

**Regions of High SV**

The soybean CNV data showed elevated SV within clusters of NB- and RLP-encoding genes. These are common classes of disease resistance genes (R genes) that have been shown to frequently have functions in pathogen perception and signaling of plant host defense responses (Krujit et al., 2005; DeYoung and Innes, 2006). Genes involved in immunity, environmental response, and defense have also been reported to be enriched within CNV regions in human and other mammalian genomes (Feuk et al., 2006; Nguyen et al., 2006; Perry et al., 2006; Nicholas et al., 2009; Liu et al., 2010; Gokcumen et al., 2011; Hou et al., 2012). The co-occurrence of CNV with clusters of NB- and RLP-encoding genes in soybean is intriguing when one considers the biological function of the loci, the potential mechanisms for acquiring the diversity necessary for recognizing new pathogens, and the consequences possibly imposed upon the genome by breeder-assisted positive selection.

It has been proposed that the plant’s first line of defense is through the perception of highly conserved pathogen-associated molecular patterns (PAMPs) by cell surface pattern recognition receptors (Jones and Dangl, 2006; Zipfel, 2009). This perception can lead to a non-race-specific resistance termed PAMP-triggered immunity. To defeat or suppress PAMP-triggered immunity, pathogens have evolved effector proteins. In turn, plants have evolved R genes that act to directly or indirectly perceive the pathogen effector protein and signal a defense response (Jones and Dangl, 2006). Unlike the PAMP/pattern recognition receptor relationship, which may be highly conserved (Boller and Felix, 2009), the effector/R gene relationship exists in flux and results in race-specific resistance as the pathogen evolves to escape perception and R genes coevolve to adapt to the evolved pathogen (de Wit et al., 2002; Takken and Rep, 2010; Ravensdale et al., 2011). Therefore, the importance of any given R gene may change depending on environmental circumstances. The R gene may be essential for survival in the presence of a pathogen harboring the cognate effector protein. However, in the absence of the pathogen, the R gene may become dispensable.

Structural changes, particularly gene loss or gene gain, may occur within R gene clusters as a consequence of natural and/or artificial selection. Unequal crossing over within existing gene parts or recombination between diverged haplotypes may give rise to new R genes with novel functions, some of which may be beneficial in combating newly arisen pathogen strains. Thus, there may be a gain in fitness for the creation of an R gene with a new specificity. Conversely, in the absence of a pathogen that can be recognized by a given R gene, there is no fitness cost to the loss of the unutilized R gene. In fact, there may be a fitness cost to maintaining R genes in the absence of a pathogen harboring the cognate effector protein (Tian et al., 2003; Bomblies and Weigel, 2007), driving selection to favor recombination and unequal sequence-exchange events that purge R gene copies that are no longer needed in a given environment. These factors lend themselves to the rapid evolution of R genes (Mondragon-Palomino and Gaut, 2005) and predict that R gene clusters may be hotspots for SV. Indeed, rapid evolution of NB-LRR-encoding genes was observed within homeologous regions of the *Rpg1b* locus of soybean and related species (Ashfield et al., 2012). NB-LRR-encoding genes in these regions experienced a higher rate of duplication and deletion than non-NB-LRR-encoding genes interspersed within the cluster. One might expect that other gene classes that tend to be “environmentally specific” or “conditionally necessary” may also be hotspots for SV, although perhaps undetectable with our current sample size.

The findings of this study are consistent with previous findings in soybean and other plant species. Resequencing of 80 Arabidopsis genomes revealed predicted PAV in 33% of the NB-encoding genes, 2.6-fold greater than the genome average (Guo et al., 2011). A recent CGH study identified 20 NB-encoding genes that exhibited CNV between two rice cultivars (Yu et al., 2011). An association between CNV and disease resistance genes has also been reported in a recent resequencing study of 50 rice accessions (Xu et al., 2012). Resequencing and CGH studies in maize identified a total of 20 NB-encoding genes that exhibited gene content variation among 24 different inbred lines (Springer et al., 2009; Lai et al., 2010; Swanson-Wagner et al., 2010), leading the authors to speculate that these genes may be involved in strain-specific disease resistance (Lai et al., 2010). A membrane array study conducted on multiple accessions within *Oryza*, *Glycine*, and *Gossypium* genera found that the number of NB-encoding genes varied widely both within and among the respective species (Zhang et al., 2010). Resequencing studies in maize and soybean each reported an enrichment of amino acid substitutions with a predicted large effect within NB-encoding genes (Lai et al., 2010; Lam et al., 2010). In concordance with this observation, a genome-wide analysis of segmentally duplicated NB-encoding genes in soybean reported that these genes are evolving at a higher evolutionary rate than other genes (Zhang et al., 2011).

**Prospects for Cloning Soybean R Genes**

The enrichment of CNV within NB and RLP gene clusters may complicate the detection of direct
orthologous relationships between these gene family members, even within intraspecific comparisons. The variation in gene content has often necessitated the construction of bacterial artificial chromosome libraries specific to the resistant accession in order to clone specific \( R \) genes (Ashfield et al., 2003, 2004; Bhattacharyya et al., 2005), even from species with a sequenced reference genome (Ashikawa et al., 2008; Lee et al., 2009). A number of previous studies on specific resistance loci have shown that the genes conferring disease resistance are often completely absent in susceptible genotypes. In soybean, a study of the \( Rps4 \) gene for resistance to \( Phytophthora sojae \) reported gene content variation associated with disease resistance; a specific gene deletion was associated with susceptibility in the Williams-derived genotype L89-1581 (Sandhu et al., 2004). Similarly, analysis of recombinant haplotypes within the cluster of NB-encoding genes at the \( Rsv1 \) locus for resistance to \( Soybean mosaic virus \) revealed that distinct resistant and susceptible interactions were associated with the presence or absence of the members of this cluster (Hayes et al., 2004). Additionally, bacterial artificial chromosome-based comparative sequencing of candidate \( R \) gene regions in soybean have identified several regions in which the content of NB-encoding gene clusters is highly dynamic, including many gene models that exhibit PAV among accessions (M. Graham, personal communication). The dynamic nature of NB- and RLP-encoding gene clusters decreases the utility of a single reference genome sequence for the identification and cloning of resistance genes. Diagnostic platforms capable of assessing genome-wide gene content variation among the wider soybean germplasm (beyond the Williams 82 genome sequence) may be a valuable tool for identifying candidate \( R \) genes in the future.

CONCLUSION

This report provides, to our knowledge, the first genome-wide analysis of soybean copy number and PAV among a limited sample of accessions. The CNV dynamics along the individual chromosomes were described, providing insight into the regions and chromosomes with relatively high or low rates of SV. A notable enrichment of significant CNV was identified within known \( R \) gene clusters. Furthermore, this study provides the groundwork for a deeper sampling of the germplasm that will allow for a more thorough assessment of soybean SV within the context of population and evolutionary genetics.

MATERIALS AND METHODS

Plant Material and Nucleic Acid Extraction

Seeds for soybean (\( Glycine max \)) ‘Williams 82’ were obtained from Dr. Randy Shoemaker at Iowa State University. The individual used for this study was named Wm82-ISU-01. Seeds for accessions Archer, Minsoy, and Noir 1 were obtained from the U.S. Department of Agriculture Soybean Germplasm Collection in Urbana, Illinois. The individuals used for this study were named Archer-SGC-01, Minsoy-SGC-01, and Noir 1-SGC-01, respectively. These individuals are referred to by their abbreviated or cultivar names (Archer, Minsoy, Noir 1, and Wm82) for simplicity.

Seeds were planted in individual 4-inch pots containing a 50:50 mix of sterilized soil and Metro Mix and grown under standard greenhouse conditions. Young trifoliate leaves from 3-week-old plants were harvested and immediately frozen in liquid nitrogen. Frozen leaf tissue was ground with a mortar and pestle in liquid nitrogen. DNA was extracted using the Qiagen Plant DNAeasy Mini Kit according to the manufacturer’s protocol. DNA was quantified on a NanoDrop spectrophotometer. These DNA samples were used for SNP genotyping, CGH, and exome-resequencing analyses.

Four more soybean accessions were used for additional CGH experimentation. Seeds for accessions Kingwa and Williams were obtained from the Soybean Germplasm Collection in Urbana, Illinois. Seeds for accessions Richland and M92-220 were obtained from Dr. James Orf at the University of Minnesota. Plants for these accessions were grown and DNA was extracted as described above.

Illumina Infinium Genotyping

The Illumina Infinium Select SoySNP50 chip (Q. Song, C.V. Quigley, G. Jia, P.B. Cregan, and D.L. Hyten, unpublished data) was used to obtain genotyping data for the three individual plants: Archer, Minsoy, and Noir 1. The Wm82-ISU-01 Infinium genotyping data from a previous study (Hau et al., 2011) were used for comparisons. SNP calls were made with Illumina GenomeStudioV2010.2 software. Heterozygous, ambiguous, or otherwise uninformative data points were treated as missing data. Visual displays showing the SNP profiles for the three genotypes relative to Wm82 were generated using Spotfire DecisionSite software.

CGH

The microarray used for the CGH experiments is described in detail in a previous publication (Hau et al., 2011). Briefly, the microarray consists of 696,139 unique oligonucleotide probes ranging from 50 to 75 bp in length. The probes tile the assembled soybean genome sequence at a median interval length of 1,120 bp between adjacent probes. This platform may be ordered from Roche NimbleGen by requesting the design 091113_Gmax_R5_CGH_HX3.

CGH protocols, including DNA labeling, microarray hybridization, and scanning, were performed as described (Hau et al., 2011). Genotype Wm82 was used as the Cy5 reference in all hybridizations. Data analyses followed previously described methodologies (Hau et al., 2011). Briefly, the segMNT algorithm in the NimbleScan software (version 2.5) was used to extract the raw data and make segmentation calls. The parameters of the algorithm were as follows: minimum segment difference = 0.1, minimum segment length (number of probes) = 2, acceptance percentile = 0.99, number of permutations = 10; nonunique probes were included, and spatial correction and q spline normalization were applied. The list of resulting segments was processed to identify significant segments. Segments were significant if the log2 ratio mean of the probes within the segment was beyond the threshold level for that genotype. The upper threshold was the log2 ratio value of the 95th percentile of all data points for each individual genotype. The lower threshold was the reciprocal negative value of the upper threshold. The thresholds for Archer, Minsoy, and Noir 1 were \( \pm 0.437, \pm 0.449, \) and \( \pm 0.421, \) respectively. Following manual inspection, each segment was compared with the soybean high-confidence gene list to identify genes and repetitive elements within significant segments.

To compare the relative SV among Archer, Minsoy, and Noir 1, we calculated the variance between the Archer-Wm82, Minsoy-Wm82, and Noir 1-Wm82 hybridization log2 ratios for each probe on the microarray. The mean variance along a sliding window of 250 probes was calculated and plotted for each chromosome. Visual displays of the CGH data with respect to the significant CNV probes and PAV genes were generated using Spotfire DecisionSite software.

Additional CGH experiments using genotypes Kingwa, Williams, Richland, and M92-220 were also performed as described above.

Exome-Resequencing and Data Analysis

Exon DNA was captured from the Archer, Minsoy, Noir 1, and Wm82 samples using the NimbleGen soybean exome chip (design 100310_Gmax_public_exome_cap_HX3) and was resequenced using the Illumina Genome
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Analyzer II platform. The exome capture and 76-bp paired-end sequencing were performed as described previously (Hau et al., 2011).

Sequence reads were aligned to the soybean reference genome sequence (Schmutz et al., 2010) using SOAP2 (Li et al., 2009), as described previously (Hau et al., 2011). The number of reads per gene model exon was calculated as described (Hau et al., 2011). The total number of reads mapping to exons for the Wm82 sample was 29,054,888. The respective numbers of reads mapping to exons for the Archer, Minsoy, and Noir 1 samples were 25,915,052, 19,878,546, and 22,480,515, respectively.

The exon read count for each Glyma gene model was used to detect gene content differences among the Wm82, Archer, Minsoy, and Noir 1 samples. Read counts were normalized by applying a correction factor to each sample that adjusted the total number of read counts to 19,878,546 (the number of Minsoy read counts). Gene content variants were defined as any gene model that had a minimum of 30 read counts in at least one genotype and zero read counts in at least one genotype; 133 such gene models were identified. To subclassify the gene content profiles, a cutoff of six reads was set as the standard for calling a gene as “present” or “absent” within a genotype. Genotypes with six or more reads were considered present for the gene; genotypes with five or fewer reads were considered absent. Based on these presence-absence calls, the 133 gene models were subclassified into 11 different groups according to their presence-absence profile among genotypes.

To compare the exome- sequencing read counts of the 133 PAV genes with the CGH data, we computed the log, ratio for each gene in the four genotypes. Counts of zero were converted to a value of 1 to allow for the calculation of count ratios between the genotypes. Calculations and statistical analyses of the exome-sequencing and CGH data log, ratios were performed using Excel software.

PCR Validation of PAV

A subset of genes exhibiting CNV and PAV among the four genotypes were examined for PAV among a diverse set of 31 soybean accessions. PCR primers were designed for 13 soybean gene models based on genomic DNA sequences available at www.phytozome.net (version 7.0). Primer3 software (version 0.4.0) was used for primer design, targeting a product size range between 300 and 400 bp per gene model. A standard PCR protocol was executed using HotStar Taq DNA Polymerase (Qagen), with 36 cycles of heat denaturation at 95°C for 30 s, annealing at 55°C for 30 s, and extension at 72°C for 40 s after an initial denaturation at 95°C for 1 min. PCR products were run on 1.4 agarose gels and stained with ethidium bromide. PCR bands were visually examined for PAV among a diverse set of 31 soybean accessions. PCR primers were designed for 13 soybean gene models based on genomic DNA sequences available at www.phytozome.net (version 7.0). Primer3 software (version 0.4.0) was used for primer design, targeting a product size range between 300 and 400 bp per gene model. A standard PCR protocol was executed using HotStar Taq DNA Polymerase (Qagen), with 36 cycles of heat denaturation at 95°C for 30 s, annealing at 55°C for 30 s, and extension at 72°C for 40 s after an initial denaturation at 95°C for 1 min. PCR products were run on 1.4% agarose gels and stained with ethidium bromide. PCR bands were visually scored as either present or absent for each genotype template.

Comparison of Regions of SV with Known QTL

For comparisons of SV and known QTL, we defined the regions with the greatest amplitude of CNV variance as regions exhibiting over 0.5 variance between the log, ratios of the Archer, Minsoy, and Noir 1 hybridizations (Fig. 4). The genome sequence coordinates of markers from the soybean consensus map 4.0 (Hyten et al., 2010; available at http://www.soybase.org) were used to estimate the genomic region associated with these high-amplitude CNV regions. Genetic locations of QTL and genes for disease resistance from previous mapping studies were estimated from the soybean consensus map 4.0 (www. soybase.org).

Identification and Enrichment Analysis of Gene Classes Associated with Genomic SV

The genes associated with PAV and significant CNV regions were subjected to GO and other analyses to identify gene classes enriched within SV regions. The subset of 133 PAV gene models identified in the exome-sequencing data was selected for this analysis. Additionally, we used the CGH data to select the subsets of gene models in which any part of the predicted coding region overlapped with or resided within a significant CNV segment. All gene annotations were estimated from the longest open reading frame of the soybean 46,430 high-confidence predicted protein-coding genes (Schmutz et al., 2010). GO designations (Berardini et al., 2004) were assigned based on the highest BLASTp (Altschul et al., 1997) hit of soybean predicted peptides to the Arabidopsis (Arabidopsis thaliana) protein database with an expectation threshold of $1 \times 10^{-20}$. Soybean sequences were classified according to Arabidopsis biological process GO designations. Protein domains were predicted by Pfam, with the cutoff defined by gathering thresholds (Finn et al., 2010).

To analyze the relative effects of structural features versus selection pressures, two large protein families involved in disease resistance, the NBR and RLP families, were identified from the total soybean protein set. NF family members were defined by the presence of an NB-ARC (nucleotide-binding adaptor shared by APAF-1, R proteins, and CED-4) domain (Pfam: PF00931). RLPs were defined by the presence of the Pfam LRR domain, the conserved LRR-containing C domain, and a transmembrane domain. A hidden Markov model was built from an alignment of the C3 domain from Arabidopsis RLPs (HMMER3; Eddy, 2009). Proteins with predicted LRR and C domains, with no other predicted Pfam domain with an e-value less than 0.5 (Tor et al., 2004), and with an identifiable C-terminal transmembrane domain were categorized as RLPs. Transmembrane topology was predicted by hidden Markov models using TMHMM (Krogh et al., 2001; Kahsay et al., 2005). Additionally, RLP-like proteins were identified as gene models with the following characteristics: bidirectional best BLASTp (threshold $1 \times 10^{-20}$) hits to known functional RLPs or RLPs characterized in Arabidopsis (Tor et al., 2004), containing a predicted LRR and C domain, and not containing other predicted Pfam domains (but lacking a predicted C-terminal transmembrane domain). The RLP and RLP-like gene models were grouped together into an “RLP encoding” family for downstream analyses.

Enrichment or depletion of a GO category or protein domain was determined by a hypergeometric distribution (Fisher’s exact test) with adjustment for multiple hypothesis testing achieved by resampling methods implemented by the FuncAssociate 2.0 program using 10,000 simulations (Berriz et al., 2009). All genes eligible to be called within a CNV region or as a PA V were used as reference, 46,275 and 43,530 genes, respectively. Adjusted P values were double-doubled to account for the two-sided Fisher’s exact test.

Gene families were gathered via BLASTCLUST (Altschul et al., 1997) with greater than 50% amino acid identity over more than 70% of the sequence length. Gene families were defined as large (more than 10 members) or small (two to 10 members). Based on their genomic distribution, gene family members were categorized as isolated or clustered. A cluster was defined as two or more members of a family with a maximum of eight intervening genes (Richly et al., 2002; Meyers et al., 2003). Tandem repeats within the genomic sequence of individual genes were predicted de novo with the Tandem Repeat Finder (Benson, 1999). Settings were modified from defaults to include a maximum repeat period of 2 kb, and repeats were filtered to a minimum length of 30 bp. Tandem repeats and gene family membership and distribution are available in Supplemental Tables S9 and S10. The coordinates of TEs were downloaded from SoyTEdb (Du et al., 2010). For each gene, it was determined whether the gene start or end coordinates were within 5 kb of the start or end coordinates of a TE.

All CGH data from this study are freely available from the Gene Expression Omnibus database (http://www.ncbi.nlm.nih.gov/; accession no. GSE28905). All exome-sequencing data are freely available on the National Center for Biotechnology Information Short Read Archive database (http://www.ncbi.nlm.nih.gov/; accession no. SRA039969).

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure S1. CNV among soybean genotypes.

Supplemental Figure S2. Frequency of shared and unique CNV associated with soybean gene models.

Supplemental Figure S3. CGH profiles among technical replications of the Minsoy-Wm82 and Noir 1-Wm82 hybridizations.

Supplemental Figure S4. Relationship between genomic SV and nucleotide SNP polymorphism among soybean genotypes.

Supplemental Figure S5. Distribution of presence-absence gene content variants among the four soybean genotypes.

Supplemental Figure S6. Distribution of presence-absence gene content variants among 31 diverse accessions.

Supplemental Figure S7. Frequency of shared and unique CNV associated with soybean gene models across five comparisons.

Supplemental Table S1. The gene models within regions of significant CNV between Wm82 and each of the three other genotypes.
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